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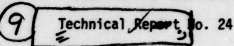






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by

10 Minoru Tsutsui

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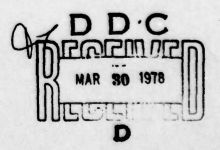
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APSTRACT (Continue on reverse side if necessary and identify by block number)

A broad review on current progress in chemistry of organometalloporphyrins is reviewed. An emphasis was placed on the development of metallocarbonyl-porphyrins, dinuclear metalloporphyrins and Skewered metalloporphyrins.



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## NEW TRENDS IN THE CHEMISTRY OF ORGANOMETALLOPORPHYRINS

Minoru Tsutsui

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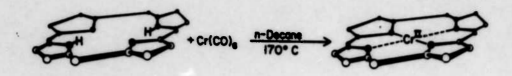
Organometalloporphyrins in general are defined as synthetic metalloporphyrin complexes containing at least one direct metal-carbon bond. However, the coenzyme fo vitamin  $B_{12}$  (Figure 1) containing an adenosyl group linked to cobalt by a direct cobalt-carbon  $\sigma$ -bond is the

first example of a naturally occurring organometallocorrin complex (similar to organometalloporphyrins in molecular structure) and the first-known stable organocobalt deriva-The determination of tive. the structure of the vitamin B, coencype by x-ray crystal-lography prompted several studies of the synthesis and properties of cobalt-alkyl compounds. Johnson and his co-workers extended their extended their studies to the porphyrin series by reaction of pyridinobromocobalt(III)actioporphyrin-I with a variety of alkyl- and aryl-magnesium halides in anhydrous 1,2-dimethoxyethane (DME) to give the corresponding elkyl- and apyl-cobalt(III) derivatives. Ethyl and pderivatives. tolyl-iron(III) derivatives? of aetioporphyrin-I were also prepared by a similar method. The first-row transition metal. derivatives of organometalloporphyrins were thus synthesized for the first time. In an alternative synthesis, Johnson and his co-workers were also able to prepare the identical alkyl cobalt(III)actioporphyrin-I complexes as that mentioned above, by reduction of cobalt (II)actioporphyrin-I with 1% sodium emalgam in DME to form cobalt(I) species, which then react with alkyl halides to give the final products. This method was also used for the preparation of hydroxyalkyl-

SPATIAL STRUCTURE OF COENZYME B<sub>12</sub>

cobalt(III)actioporphyrin-I complexes. Both the cobalt(III) and the iron(III) organometalloporphyrins are sensitive to complexes. Both the cobalt(III) and decompose by homolytic fission of the metal-carbon bond.

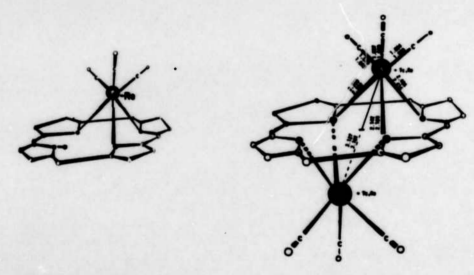
The use of metal carbonyls for the insertion of metal ions into porphyrins was first introduced by Tsustui and his co-workers in 1966(Figure 2). This method is probably one of the most important developments in porphyrin chemistry within the last two decades. In addition to a number of previously reported metalloporphyrins, the reaction of metal carbonyls and metla carbonyl halides 12,21,24-29 with neutral porphyrins has lead to the synthesis of new metalloporphyrin complexes of chromium, molybdenum, technetium, ruthenium, rhodium, rhenium and iridium. 10-32 Except for the chromium and molybdenum complexes, carbonyl groups



USE OF Cr(CO)<sub>6</sub> FOR INSERTION OF Cr<sup>II</sup> TO PP

are retained by the metals in the new metalloporphyrin complexes.

By reaction of dirhenium decarbonyl, Re<sub>2</sub>(CO)<sub>10</sub>, or ditechnetium decacarbonyl, Tc(CO)<sub>10</sub>, with mesoporphyrin IX dimethylester, H<sub>2</sub>MPIXDME, if refluxing decalin under argon, Tsutqui<sub>3</sub>2nd his co-workers have successfully prepared two unusual rhenium organometalloporphyrins 1,32 (H-MP)Re(CO)<sub>2</sub>, I, and MP[Re(CO)<sub>3</sub>]<sub>2</sub>, II, a pair of technetium organometalloporphyrins, 33,34 (H-MP)<sub>1</sub>Tc(CO)<sub>2</sub>, III, and MP[Tc(CO)<sub>3</sub>]<sub>2</sub>, IV, and a mixed rhenium technetium organometalloporphyrin (CC)<sub>3</sub>ReMTc(CO)<sub>2</sub>, Y. A single crystal z<sub>0</sub>ray diffraction analysis of  $\mu$ -[meso-tetraphenylporphinato]bis[tricarbonylrhenium(1)], TPP[Re(CO)<sub>3</sub>]<sub>2</sub>, V, (Figure 3), has shown



Porphyrin - Tridentate Ligand

Porphyrin-Hexadentate Ligand

#### FIGURE 3

that each rhenium ion is bonded to three nitrogen atoms and that two rhenium atoms are bonded to one porphyrin on opposite sides of the plane of the porphyrin molecule.

The metal ions in these complexes, I-VI, sit out of the plane of the porphyrin molecule. The monorhenium and monotechnetium complexes, I and III, where the porphyrin noiety acts as a tridentate ligand, resemble Fleischer's proposed "sitting-atop complex" 30,39 and are good models for the intermediates in the insertion of a metal ion into porphyrin. The dirhenium, ditechnetium, and mixed rhenium-technetium organometalloporphyrin complexes, II, IV, Y, and YI, where the porphyrin moiety acts as a hexadentate ligand, are examples of the first isolated stable home—and hetero-dinuclear organometalloporphyrin complexes. The monorhenium porphyrin complex, I, reacts with Re<sub>2</sub>(CO), or Te<sub>2</sub>(CO), in refluxing decalin to form the dirhenium porphyrin complex, II, and the mixed rhenium technetium porphyrin complex. Replacement of the pyrrolic proton (NnH) of the monorhenium porphyrin complexes. The monortechnetium porphyrin complex, III, (Figure 4), behaves in a different manner by disproportionating to form a ditechnetium porphyrin complex, IV, and the free porphyrin, R.MPIXDME, by heating in refluxing decalin. This unusual coordination phenomenon has never been reported. Such a reaction was not observed on heating monorhenium porphyrin complex, I, in refluxing decalin. It seems that both the rhenium and technetium dimetalloporphyrin complexes are are thermodynamically more stable than the monometalloporphyrin complexes, because a reverse reaction of NP[M(CO)<sub>3</sub>]<sub>2</sub> to (H-NP)-M(CO)<sub>3</sub>,

(N=Re or Tc), could not be detected between NP[N(CO)], and H\_NFIXING in refluxing decalin for either the rhenium of technetium dimetalloporphyrifi complexes.  $^{31-34}$ 

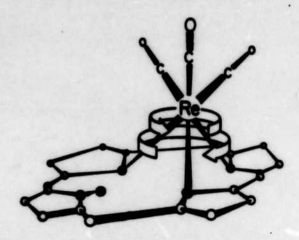
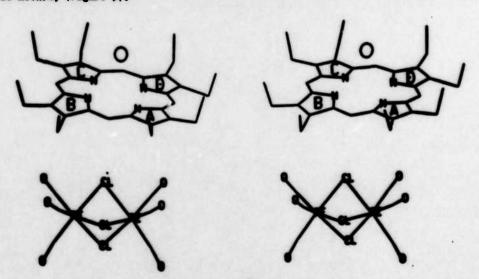


FIGURE &

Temperature-dependent <sup>1</sup>H new spectral changes for HTPTNe(CO), and HDPNe(CO), dissolved in 1,1,2,2-tetrachlorethane showed fluxional behavior of Re(CO), group. This phenomenon is best explained by the intramolecular rearrangement of the metal carbonyl group among the four ring nitrogens of porphin and also movement of the H-H; it can also be regarded as as intramolecular substitution at rhenium or pechnetium, (Figure 4).35

We have prepared a new salt type complex of porphyrins, monocation octaethylporphyrin tri--u-halogeno-hexacarbonyldirhenate(I) from the reaction of Re(CO)<sub>c</sub>Cl and H<sub>c</sub>OEP in a 2:1 mole ratio in decalin. The structure of the complex was elucidated by the x-ray diffraction enalysis method, (Figure 5).<sup>30</sup>

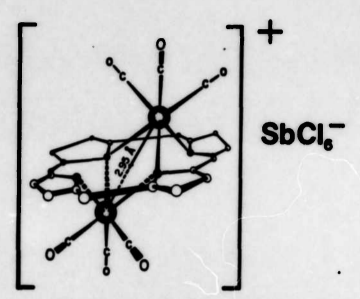


STRUCTURE OF (H30EL), [Ne<sup>5</sup>(co<sup>6</sup>)cr<sup>3</sup>]\_H<sup>5</sup>0

FIGURE 5

[Re(CO),]\_TPP has been oxidized by SbCl, in dichloromethane to yield [Re(CO),Cl],TPP · 25bCl, and [Re(CO),][Re(CO),Cl] · SbCl. An x-ray determination of the structure of these complexes provides definite evidence for a "skewered complex", that is a metalloporphytin in which a metal-metal bond exists through the "hole" of the macrocycle, (Figure 6).

Two different methods were employed by Fleischer and his co-workers in preparing the rhodium and iridium porphyrin complexes. In one, the freshly prepared metal carbonyl halides,  $[Rh(CO)_{c}Cl]_{c}$  and  $[Ir(CO)_{c}Cl]_{c}$ , were allowed to react with the porphyrins in glacial acetic acid solution to form the respective metalloporphyrins, (Figure 7). In the second method,



First Skewered Complex (Radical Metalloporphyrin)

the cyclooctene complexes of rhodium and iridium were found to be reactive intermediates useful in the metalloporphyrin formation, (Figure 8). In both sthods, incorporation of rhodium into the porphyrin was more readily achieved than was that of iridium. It is of interest that in the metalloporphyrins prepared by these methods, the iridium porphyrins retain the earbon monoxide ligand while reeding and other metals do not. The carbon oxide is very tightly bound to the iridium porphyrin moiety; heating, pumping, boiling in pyridine, or reprecipitating from concentrated sulfuric acid solution does not remove the

FIGURE 6 [r(00), CI] PH, . HOP(IX)DEEH, MP(IX)DEEH,

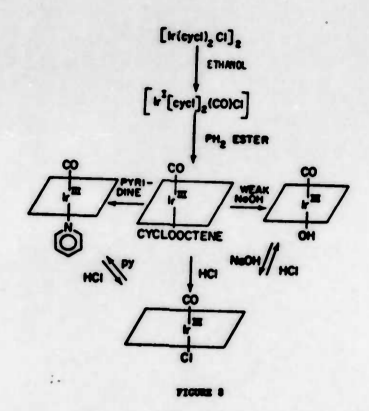
REACTION SCHEME FOR IRIDIUM PORPHYRIN PREPARED VIA Ir(CO),Cl

solution does not remove the earboayl group from the complex. 2½

By the reaction of [Rh(CO)<sub>2</sub>Cl]<sub>2</sub>

with meso-tetraphenylporphine, H<sub>2</sub>TPP, in refluxing bensene two stable organometalloporphyrin derivatives of rhodium, Rh<sup>11</sup>CO(TPP) · Cl and (σ-phenyl)Rh<sup>12</sup>-(TPP) · Cl, were separated by chromatography on an alumina column by Fleisher and his co-workers<sup>21,20</sup>.

Techida and his co-workers agragable to prepare two novel dinuclear rhodium(I) organo-metalloperphyria complexes III and VIII, by modifying Fleischer's reaction conditions for the appearation of Rh CO(TPF) · Cl and (s-phenyl)Rh (TPP) · Cl in refluxing benfor the meparation of Rh "CO(TPP) • Cl and (s-phenyl)Rh" (TPP) • Cl in refluxing bensene. "Ctaethylporphyrin, OEPH, or (N-methyl) octaethylporphyrin reacts with
[h(CO)\_Cl], in bensene solution at from temperature under nitrogen atmosphere to produce
and fitt. From the spectral data and the experimentally determined molecular weights,
was formulated as an acid, H [OEP • Rh\_(CO) Cl], which contains a Rh-Cl-Rh bridge.
The proton num and infrared spectral data indicate that the [Rh(CO)\_Cl], moiety is maintained and the N-H and N-CH, bonds exist in VII. Since the Rh-Rh distance in [Rh(CO)\_Cl],
has been reported to be 3.12 A, and the distance between the two adjacent nitrogen atoms of
planar porphyrin is about 2.9 A, it was assumed that the two Rh atoms of the [Rh(CO)\_Cl],
moiety are bonded to the two adjacent nitrogen atoms of the porphyrinato core of VIII, as



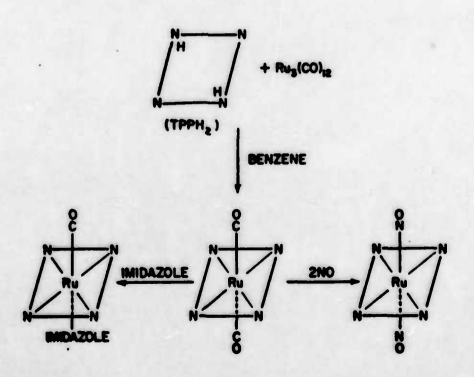


FIGURE 1

shown in Figure 9. Upon exposure to air XII was slowly oxidized to form a rhodium(III) chloro complex of octaethylporphyria, Rh (I(OZP) · 2H\_O, which can further react with alkyl lithium to give an alkyl-rhodium complex, another example of rhodium organometalloporphyrin complex. However, YIII behaves in a different manner to give the identical alkyl-rhodium complex either by gentle heating in chloroform of chromatography on silica gell (Pigure 9). This phenomenon of alkyl migration from a nitrogen atom to a metal ion is reported for the first time. The alkyl migration may proceed concertedly with oxidation of rhodium(I) to rhodium(III). The H-CH, bond fission seems to be facilitated by the aid of e low-valent rhodium ion. The reaction of (H-ethyl)octaethylporphyrin with [Rh(CO)\_Cl], yields a rhodium(1) complex similar to YIII, which is also easily oxidized to CH\_CH\_ 1 oxidation of metal oxidation and alkyl migration is still unknown.

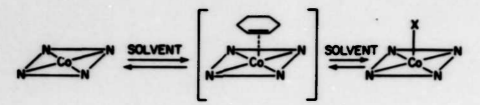
Both rethenium carbonyl, Ru<sub>3</sub>(CO)<sub>12</sub>, and ruthenium carbonyl halide, [Ru(CO)<sub>2</sub>Cl<sub>2</sub>]<sub>2</sub>, react with tetraphenylporphing to give the Identical product, monocarbonyl-ruthenium(II) tetraphenyl porphine (II) tetraphenyl porphine (II) tetraphenyl porphine (III) tetraphenyl tetraphenyl tetraphenyl (III) tetraphenyl (III) porphyrin photodimer with carbonyl tetraphenyl (III) porphyrin photodimer with carbonyl tetraphenyl (III) porphyrin photodimer with a metal-metal bond. (It is of interest that MFRuCO in bensene solution reacts apporthly with excess mitric oxide to form a dimitrosyl-ruthenium(II) meso-porphyrin complex.

Since the successful isolation and purification of nitrogenase, the ensyme system that fixes molecular nitrogen to amonia in living organisms, many studies have been made to find a nitrogenase model. Fleicher and co-workers created a model system compasisting of the water soluble mass-tetra(p-sulfonatophenyl)porphinatocobalt(III), Co TTPS, and sodium borohydride and found it to be very effective catalyst for the reduction of the substrate of nitrogenase. For example, acetylene was reduced catalytically in this system to ethylene and ethane. A possible mechanism for this catalytic system was proposed to include cobalt-acetylene and cobalt-achylene metalloporphyrin v-complexes as intermediates (Pigure 10). For the induced oxidation of cobalt(II) mesoporphyrin IX dimethyl ester,

POSSIBLE MECHANISM FOR REDUCTION OF ACETYLENE BY A COBALT(III) PORPHYRIN SYSTEM.

#### FIGURE 10

CO<sup>II</sup>PINE by unsaturated hydroacrbons, an olefin-cobalt w-complex intermediate was proposed (Figure 11), Recently, the formation of an 1:1 adduct between cobalt(II) tetra-phonylposphyrin, Co<sup>II</sup>TPP, and carbon monoxide was reported by an esr study of a frozen



#### PROPOSED REACTION INTERMEDIATE FOR INDUCED OXIDATION OF COBALT(II) PORPHYRIN BY CYCLOHECENE

#### PIGURE 11

solution of toluene. The paramenagtic low-spin d complex of Co TPP has a single unpaired electron in the dz orbital for s bonding with carbon monoxide, which provides an opportunity for the examination by esr of o spin delocalization to carbon monoxide. Similarly, the reversible binding of carbon monoxide to iron(II) protonorphyrin II in piperidine was identified by its ear spectrum in frozen solution.

The o-interaction in carbon monoxide to metalloporphyrin adducts and the proposed olefinmetal w-complex intermediates for induced oxidation of Cobelt(II) porphyrin by unsaturated hydrocarbon and the catalytic reduction of acetylene in Co TPPS and MaBH, model system has brough new examples of formation of organometalloporphyrins.

ACCOUNTEDGEMENT-This research was supported in part under a contract of the Office of Maval Research.

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